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P. 02

ATTACHMENT A

PATENT

#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

or. Pu

HIV IMMUNOASSAYS USING BYNTHETIC ENVELOPE POLYPEPTIDES (AS AMENDED)

#### DECLARATION

Honorable Commissioner of Patents and Trademarks Washington, D.C. 20231

Sir:

- I, John A.T. Young, do hereby declare as follows:
- 1. I received my Ph.D. in Human Genetics from Imperial Cencer Research Fund and University College, London, United Kingdom in 1987 having previously received a B.S. in Biochemistry from the University of Dundee in 1983.
- 2. I am currently an Assistant Professor, Department of Microbiology and Molecular Genetics, Harvard Medical School. My Curriculum Vitae is attached as Exhibit
- 3. I have read and understand Luciw at all, application Sarial No. 08/089,407 and Luciw at all application Sarial No. 08/667,601 (\*501) as well as the Office Action mailed January 23, 1896.

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- 4. One of ordinary skill in the art in 1994 understood the term "symmetic peptide" to mean a peptide prepared by chemical symthesis. The term "symthetic" was used to describe a paptide synthesized by chemical means in numerous publications prior to the October 31, 1984 filing date of perent application Serial No. 09/087,501. Representative publications (there are still others) include Akman 1984, Barkes 1994, Balks 1984, Date 1983, Green 1983, Hintz 1983, Hirayama 1982, Jacob 1983, Jolivet 1983, Lieu 1978, Morrow 1983, Morrow 1984, Mulliar 1983, Pacalla 1983, Rothbard 1984, Rougen 1984, Shervood 1983, Shi 1984, Sutcliffe 1983, Tamura 1982, and Wabuke-Bunod 1984." The articles were published in a variety of well-known journals, including those read by a general soluntific audience (e.g., PNAS and Science) as well as those read mainly by virologists and immunologists (e.g., Journal of Virology and Molecular Immunology). These are the Journals that one skilled in the art would be expected to review.
- 6. Poliziving 1984, the term "synthetic" was still understood by those skilled in the art to make a populae synthetized by chemical means. This is illustrated by the following sentence taken from Chapter 5 under the sub-heading "Synthetic paptides" of a widely-circulated laboratory research manual (Harlow, E., and D. Lane. 1988, Antibodies: a laboratory manual. Cold Spring Harbor Laboratory. Cold Spring Harbor, N.Y.): "Populaes are normally synthesized using the solid-phase techniques phoneered by Mamfield (1983)." The term is still so-understood today.

The full citation for each of the references cited in this declaration is included in Exhibit 2.

- P. 64
- 6. The prior ort was capable of making a clear distinction between a symmetric peoplide (i.e. one symmetrical means) and a peoplide fragment generated by some other means. See, Date 1983, Highwarms 1982, Lieu 1975, Morrow 1983, Rothbard 1984, and Sharwood 1983.
- 7. Prior to October 31, 1884 one skilled in the art was fully capable of synthesizing peptides of considerable length. Specific examples of synthetic polypeptides containing as many as 40 amino acids were reported in the art prior to October 31, 1884. Ten of the above-mentioned articles (Altman 1884, Baricas 1884, Dale 1889, Hirsystem 1882, Jacob 1883, Multer 1883, Rombard 1884, Shi 1884, and Wabuke-Sunoti 1884) report synthetic pagitides (i.a.! peptides made by charmost synthesia) having lengths of from 15 to 24 amino acids and one article (Sellet 1984) reports a 37 amino acid synthetic papides. Raid (1981) employed a 34 amino acid synthetic paptide.
- 8. Immunoacceye employing synthetic peptides such as claimed in the subject application were known in the art in 1984. Those techniques included ELISA analyses which employed peptides immobilized on microtion plates, test sera, and emayme-coupled secondary ambitates (e.g. Altman 1984, Beliet 1984, Green 1983, Jolivet 1983, Rothbard 1984, Wabuka-Bunoti 1984). Those techniques also included solid-phase redigimmunoacceya that employed immobilized synthetic peptides, test sera, and 1864, labeled protein A (Jecob 1983, Morrow 1984, Pacella 1983, Rothbard 1984,). Other medicals were also known in the art in 1984 for detecting specific imatections between synthetic peptides and ambitation including radioimmunoacceys that employed

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radioactively-labeled peptides or amibodies (e.g. Barlas 1984, Hintz 1982, Rougon 1984, Shi 1884, Tamura 1982).

- 9. The statement at page 3 of the '501 specification that 'synthetic peptides may also be prepared' would have been understood by one of ordinary skill in the art in October 1984 as a reaching that such synthetic peoples would be used in the immunosessys described in the '801 specification. The '501 specification at pages 11, 14 and 15 specifically teaches that one use for the polypeptides of the invention is as amigens in a variety of immunosessys. One skilled in the art would not infer from the teaching of the patient specification that production of synthetic peptides would be a teaching of a variety at the specification that would be lad to use the synthetic peptides in immunosessys just as the specification teaches.
- 10. The HIV nucleotide and antino acid sequences provided in the '501 parent application enabled one of ordinary skill in the art in October 1984 to identify synthetic HIV antigenic peptides, i.e., peptides containing an instrumogenic amino acid sequence. To demonstrate this, I performed a hydrophilicity analysis of the ARV-2 Env sequence, according to the Hopp protocol (Hopp 1981, Hopp 1983). The directions in Hopp, together with the hydrophilicity values given in Hopp 1981, permit a streightforward analysis that was eatily within the skill of the err in October 1984. The confirmation of antigenicity was also within the skill of the err in October 1984. The confirmation of antigenicity was also within the skill of the err in 1984. An arrigen could be accorded by using it in an immunoassay such as the prior art immunoassays identified in Peregraph 8 and testing it with sens of patients known to be infected. This acreening process is the technique that is, in fact, disclosed in the Hopp references.

- 11. Employing the Hopp protocol, the most hydrophilic region of ARV-2 2m, was identified as residues 738-743 (ERDRDR). Synthetic peptides derived from HIV Env that contain thisse amino acid residues are recognized by a proportion of AIDS patient entisera as demonstrated by later actual trass. (Broliden 1992, Goudarnist 1980, Karnedy 1986). The second-most hydrophilic region was identified as residues 653-658 (ERNECE). Synthetic peptides containing this region of HIV Env are also recognized by sara from HIV infected Individuals (Broliden 1982, Goudarnit 1990, Krowka 1991). The third most hydrophilic region of ARV-2 Env, residues 739-738 (EEEGGE), overlaps the first hydrophilic region. Synthetic peptides committing this third region of HIV Env are recognized by sara from HIV Infected individuals. (Broliden 1982, Goudarnist 1990, Karrascy 1983). The region containing residues 505-510 (QREKRA) was also identified as being highly hydrophilic. This finding was noted using the same computer analysis by Pauletti (1985). Synthetic peptides derived from HIV Env containing all or most of these residues are recognized by AIDS patient analysis (Broliden 1992, Kennedy 1987, Knowka 1991, Manhoheysikova 1993, Palear 1987, Stredard 1992).
- 12. Employing the Hopp protocol, the most hydrophilic region of ARV-2 Geg, was identified as residues 102-107 (ERREE). Synthetic peptides derived from HIV Geg that combin these amino acid residues are recognized by a proportion of AIDS patient antisera as demonstrated by fater actual tests. (Jiang 1992). The second-most hydrophilic region was identified as residues 109-114 (NKSKIGG). Synthesic paptides containing this region of HIV Geg are immunoganic and are recognized by eare from HIV infected individuals (Jiang 1992).

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- 13. The HIV sequences provided in the '801 parent application also enabled one of ordinary skill in the ert in October, 1984 to identify antigenic HIV Env linear epitopes by still other textiriques. One other approach known in the ent, was to generate one or a panel of several synthetic peptides derived from the polypeptide sequence and test each peptide for antibody reactivity. The generation of one or a panel of synthetic polypeptides from a single protein was a routine matter in 1984.
- 14. A penal of sight population teach 13-15 aminu acids in length) of interleukin-2 was generated by Altman (Altman 1994) and a panal of five synthetic paptides (8 to 16 amino acids long) derived from adenovirus 1915 and 6916 proteins was generated by Green 1983). In addition, Sutsiffic generated a panel of 12 paptides from MuLV polymerase generated a panel of 18 paptides from the rabbes glycoprotein gene. (Sutcliffe 1983)
- 15. Prior to October 1984, those skilled in the art know that a proportion of antibodies raised against rective promitte could recognize epitopes contained on synthetic paptides derived from a protein sequence (Pembarit 1984; Leach 1983) or contained on proteolytic protein fragments (Lando 1982).
- 16. Based on the information described herein, those sidiled in the art could have, without undue experimentation, used the sequence of ARV-2 Env provided in the 1601 application to generate synthetic peptides representing most of the HIV glycoprotein. These peptides could then have been tagged using standard assays known in the art, and imminispenic regions of HIV Env identified.

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- 17. I have reviewed in detail Montagnier, Solonce, 225, 63-66 (July, 1984) and Schupbach et al., Science, 224, 508-505 (May, 1984). In my opinion these articles would not have enabled one skilled in the art to prepare a synthetic HIV envelope polypeptide sequence for use in an immunoassity without undue experimentation. I conclude this for the following ressaus:
  - These articles did not provide any MIV nucleotide or arring acid sequence information.
  - Although HIV proteins were purportedly identified by immunoblorting in these publications, a person of ordinary skill in the art would not have been able to produce sufficient quantities of any of these viral proteins for sequencing. Sufficient quantities could not have been produced because cultures of primary human cells falled to produce significant quantities of HIV, as the virus is cytopathic and repidly killed the infected virus-producing cells. Therefore, a person of ordinary skill in the art, attempting to generate sufficient quantities of HIV proteins for detailed characterization, would have I) had to obtain an appropriate established cell into known to produce HIV and it) had to have a knowledge of the practice conditions required for infecting these cells and for malmostring the infected cells for long periods of time in culture.
  - o) By October 31, 1984, the Gallo and Montagnier groups had reported cell lines that could be used to produce significant levels of MIV (Popovic 1984, Montagnier 1984). Gallo and Montagnier were world

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the of "ordinary skill in the ert". At the time of the '601 application date, the precise origin of the cell line used by the Galle group had not been disclosed (Popovic 1884). The Montagnier group used cells generated by fusion between HIV producing primary T cells and EBV-transformed B-cells (Mortagnier 1984). It would not have been possible for a scientist of ordinary skill in the art to have used the same technique to produce cell lines that were identical to those described by the Mortagnier group. Even if a scientist of ordinary skill in the art had attempted to obtain the cells described by the Galle and Montagnier groups, I am not aware of any evidence that these cell these were being distributed freely to the public at the time of the '501 application date. Furthermore, the precise culture conditions required for maintaining HIV-infected cells in culture had not been disclosed.

18. The ennouncement by the Gallo group that HTLV-III was related to HTLV-I and II, such as contained in Gallo et al. (1983) and Arya et al. (1984), led workers such as Crang to incorrectly presume that the Env gene was located at the same position in the HIV and HTLV-I and II genomes. Furthermore, the Gallo group proposed that the HIV genome contains a pX or LOR region similar to those found in HTLV-I and II. In fact, as the '501 application correctly disclosed, e) HIV is not closely related to HTLVs, b) the Env gene is not closely related at the same position in the HIV and HTLV genomes and c) there is no pX or LOR region in the HIV genomes.

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- 19. The presumption that HIV was closely related to HTLV-I and II led the Gallo group to seriously misidentify HIV savelage proteins:
  - The Gallo group described a 65 kD HIV protein as "enveloperelated" apparently because it migrated on 808-polyacrylamide gals at a
    position similar to that of the 62-65 kD HTLV precursor envelope protein
    (Schupbech 1884). The HIV precursor envelope protein is, however, a
    180kD protein (designated gp160), a fact that only came to light after the
    "501 application fling date.
  - b) The Gallo group described a 41 kD HIV protein as "the presumed envelope antigen of the virus" (Barngedheran 1984). The 41 kD protein was shown to be an antigenic viral enructural protein (Sarngedheran 1984). However, the inescapable conclusion from this manuscript was that these workers presumed that this viral protein was envelope related because it was similar in size to the 48kD HTLV envelope protein (gp46; Sarngedheran 1984) i.e., the HIV p41 protein was equivalent to HTLV gp48. In fact these proteins are not equivalent for the following reasons:

However, these proteins are structurally distinct and perform different functions duting viral entry; the SU protein is primarily involved in receptor binding, whereas the TM protein combine the transmembrane region that anchors the envelope proteins on the virus surface. The TM protein is primarily involved at a step of viral entry following receptor binding.

- The Strand TM proteins of HIV are designated gp120 and gp41 (the 41 kD protein described by Sarngadharan 1984), respectively. The SU (gp120) protein of HIV was not described prior to the '801 emplication filing date.
- in) The BU protein of the HTLVs is gp46 and the TM protein of the HTLVs is p20E a 20 kD protein.
- 20. I have also reviewed in detail Chang U.S. application Serial No. 659,339 filed October 10, 1984 including the perial DNA sequence of Figure 3. The Chang specification (1) incorrectly describes the location of the Env gene in the HIV genome, and (2) misrepresents the sequence of the Env gene which is purported to be encorrected (i.e. wholly-contained) within the DNA sequence shown in Figure 3. An individual skilled in the entrempting to Identify the HIV Env open reading frame found in the sequence of Figure 3 would have been unable to do so.
- 21. Although Chang represents that the Figure 3 sequence 'encompasses the env region' (p. 5, lines 1-2), that is incorrect. In fact, the Figure 3 sequence commins a

partian of the pal gene, the sar gene and only approximately one-third of the envelope gene.

- 22. Moreover, the Chang Figure 3 sequence includes an error. The Figure 3 sequence includes an agra mucleatide ("A") at position 2497, a residue which does not actually exist in the HIV envelope gare. This mistake leads to a +1 translational frameshift at this position in the partial sequence of the envelope open reading frame. As a consequence of this error, this open reading frame is only correct over the region encoding the first 63 amino ecids of Env (tooluding the N-terminal algnal peptide which is removed during protein biosynthesis). The open reading frame of the Figure 3 sequence than continues with three amino acids encoded by an incorrect reading frame followed by a stop codon.
- 23. Eased on Figures 1 and 2 of Chang, a adaptist would have been completely missed about the piecement of the envelope gaps relative to restriction enzyme sites in the HIV generits, e.g., an EcoR1 site that is actually located upstream of the envelope game is shown in the Chang '339 application both as contained within the envelope game (Figure 1) and Upstream of the envelope game (Figure 2). Also, a Bigl II site, which is actually located in the envelope game, is shown in the Chang application as within the 'px' region, a region which does not exact in the HIV gamome. HIV is not closely related to HTLV-I and II, and unlike these other human removinuses HIV certainly does not contain a pX region.

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24. Based, inter also, on the above-identified defects. Chang did not enable one ciciled in the art in October. 1984 to grow, leaded and/or sequence the envelope game of HIV.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with that knowledge that willful false statements and the like on made are purchable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

nove: March 19 1997

MAR-19-1997 12:58 FRUM BANNER

D.

John A.T. Young

#### **CURRICULUM VITAE**

#### PERSONAL

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Birthdate: February 28, 1961 Kirkcaldy, Fife, Scotland Crizenship: United Kingdom

Marital Status: Married to Dr. Caroline Alexander

Social Security Number: 603-26-2953

## ACADEMIC APPOINTMENTS

1992 to 1995 Assistant Professor
Department of Microbiology and Immunology
University of California, San Francisco

San Francisco, California

1992 to 1995 Assistant Investigator
Gladstone Institute of Virology and Immunology
San Francisco General Hospital
San Prancisco, California

1992 to 1995 Member, Program in Biological Sciences (PIBS)
Cell Biology Program
University of California, San Francisco

1992 to 1995 Member, Biomedical Sciences Program University of California, San Francisco

1995 to Assistant Professor

Department of Microbiology and Molecular Genetics
Harvard Medical School

1995 to Member, Biological and Biomedical Sciences Program, Harvard Medical School

1995 to Member, Committee on Virology, Harvard Medical School

Member. Board of Tutors in Biochemical Sciences, Harvard University 1995 to

# POSTDOCTORAL TRAINING

EMBO Posidoctoral Ifallow 1987-1989

Department of Microbiology and Immunology

University of California, San Francisco Advisor: Harold B. Varmus, M.D.

Arthritis Foundation Postdoctoral Fellow 1989-1992

Department of Microbiology and Immunology

University of California, San Francisco Advisor. Harold E. Værnus, M.D.

#### **EDUCATION**

1983 University of Dundee

Dundee, United Kingdom

B.Sc., Blochemistry (Piret Class Honours)

Impedal Cancer Research Fund and University College 1987

> London, United Kingdom Ph.D., Human Genetics

Theris: Expression and Polymorphism of HI.A-D Region Genes

Ph.D. Advisor, John Trowsdale, Ph.D.

## TEACHING EXPERIENCE

1992 Co-organizer

Introduction to Cell Biology course

Medicine 412, UCSP

Discussion Leader

Cell Biology Course 212, UCSF

1993 Lecturer

The Biology of AIDS

Biomedical Sciences Minisymposium, UCSF

1993 Discussion Leader

Tissue Organization and Murphogenesis course

Biomedical Sciences 210. UCSF

Discussion Leader

Molecular Biology of Animal Viruses course Microbiology 208, UCSP

1994 Lecturer, The Biology of Virus Infection course

Microbiology 208, UCSF

Lecturer. Microbiology 201, Hervard Medical School 1996

(4 lectures, 9 discussion groups)

1997 Co-director, Virology 200, Hurvard Medical School

1997 Lecturer, Virology 200, Harvard Medical School (3 lectures)

#### COMMITTEES

1992 to 1995 Member, Dean's Advisory Committee to the UCSF AIDS Clinical Research

Center

1993 to 1995 Member, Executive Committee of the UCSF Biomedical Sciences Program

1993 to 1995 Member, UCSF Student Research Committee

1996 to Member, Virology Admissions Committee, Harvard Medical School

1996 to Member, Division of Medical Sciences Curriculum Committee, Harvard Medical

School

#### TRAINEES

1992 to 1995 Kurt Zingler

Ph.D. Thesis Student

Immunology Program, UCSP

Jürgen Brojatsch, Ph.D. Postdoctoral Fellow

Carole Bélanger, Ph.D.

Postdoctoral Fellow

Fonds de la Recherche en Santé du Québec

1993 to 1995 Lynn Connolly

M.D., Ph.D. Thesis Student

Medical Scientist Training Program, UCSF

Morgan Jonkins, M.D.

Clinical Research Pellow

Universitywide AIDS Research Program

1996 to

Heather B. Adkins

Ph.D. Thesis Student

Cummittee on Virology, Harvard Medical School

1995 to

Vincent Solomon

Ph.D. Thesis Student

Biological and Biomedical Sciences, Harvard Medical School

#### **PUBLICATIONS**

- 1. Trowschile, J., Young, J.A.T., Kelly, A.P., Austin, P.J., Carson, S., Meunier, H., So, A., Ehrlich, H.A., Spielman, R.S., Rodmer, J., and Bodmer, W. (1985) Structure, sequence and polymorphism in the HLA-D region. *Immunol. Rev.* 85:135-173.
- Young, J.A.T. and Trowsciale. J. (1985) A processed pseudogene in an intron of the HLA-DPβ1 chain gene is a member of the ribosomal protein L32 gene family. Nucl. Acids Res. 13:8883-8891.
- Trowsdale, J., Austin, P., Carson, S., Kelly, A., Lamb, J., and Young, J.A.T. (1985) Cloned HLA-D genes: Characterisation and approaches to expression and analysis of function. In: Human T-cell Clones (M. Feldmann, J.R. Lamb, and J.N. Woody, eds.), The Human Press, pp 49-57.
- Bodrner, W.F., Trowsdale, J., Young, J., and Bodner, J. (1986) Gene clusters and the evolution of the major histocompatibility complex. Phil. Trans. R. Soc. Lond. 312:303-315.
- Young, J.A.T., Wilkinson, D., Bodmer, W.F., and Trowsdale, J. (1987) Sequence and evolution of HLA DR7 and HLA-DRw53-associated β chains. Proc. Natl. Acad. Sci. USA 84:4924-4933.
- 6. Bodmer, J., Bodmer, W., Heyes, J., So., A., Touks, S., Trowsdale, J., and Young, J. (1987) Identification of HLA-DP polymorphism with DPG and DPB probes and monoclonal antibodies: Correlation with primed lymphocyte typing. *Proc. Natl. Acad. Sci USA* 84:4596-4600.
- Young, J.A.T., Lindsay, J., Bodmer, J.G., and Trowadale, J. (1988) Bpitope recognition
  by an HLA-DPα chain-specific monoclonal antibody (DPI L.1) is influenced by the
  association of the DPα chain and its polymorphic DPβ chain partner. Hum. Immunol. 23:3744.
- 8. Young, J.A.T. (1988) HIV and HLA similarity. Scientific correspondence. Nature 333:215.
- 9. Young, J.A.T. and Trowsdale, J. (1990) The HLA-DNA gone is expressed as a 1.1kb mature mRNA species. *Immunugenetics* 31:386-388.
- 10. Young, J.A.T., Baies, P., Willert, K., and Varmus, H.E. (1990) Efficient incorporation of human CD4 protein Into Avian Leukoris Virus particles. Science 250:1421-1423.
- 11. Young, J.A.T., Bales, P., and Varmus, H.E. (1993) Isolation of a chicken gene that confers susceptibility to infection by subgroup A avian leukosis and sarcoma viruses. *J. VIrol.* 67:1811-1816.
- 12. Bates, P., Young, J.A.T., and Varmus, H.E. (1993) A receptor for subgroup A Rous sarcoma virus is related to the low density lipoprotein receptor. Cell 74:1043-1051.
- 13. Connolly, I., Zingler, K., and Young, J.A.T. (1994) A soluble form of a receptor for subgroup A avian leukosis and sarcoma viruses (ALSV-A) blocks infection and hinds directly to ALSV-A. J. Virol. 68:2760-2764.
- 14. Young, J.A.T., Bales, P.F., and Varmus, H.E. (1994) A protein related to the LDL receptor in a collular receptor specific for subgroup A-avian leukosis and sarcoma viruses. In:

- Receptor-mediated Virus Entry into Cells. (B. Wimmer, ed.) Cold Spring Harbor Laboratory, Cold Spring Harbor, NY.
- 15. Young, J.A.T. (1994) The replication cycle of HIV-1. In: The AIDS Knowledge Base. (Cohen, P., Sande, M., Volberding, P., et al., eds.) Little Brown, New York, NY.
- 16. Pederspiel, M.J., Rates, P., Young, J.A.T., Varmus, H.E., and Hughes, S.H. (1994) A system for tissue-specific gene targeting: Transgenic mice susceptible to subgroup A avian leukosis virus-bused retroviral vectors. *Proc. Natl. Acad. Sci. USA*. 91: 11241-11245
- 17. Bélanger, C., Zingler, K., and Young, J.A.T. (1995) Importance of cysteines in the LDLR-related domain of the ALSV-A receptor for viral entry. J. Virol. 69: 1019-1024.
- 18. Zingler, K., Bélanger, C., Peters, R., Agard, D. and Young, J.A.T. (1995) Identification and characterization of the viral interaction determinant of the ALV-A receptor. J. Virol., 69: 4261-4266
- 19. Zingler, K. and Young, J.A.1'. (1996) Residue Trp-48 of Tva is critical for viral entry but not for high-affinity binding to the SU glycoprotein of Subgroup A avian loukosis and surcoma viruses. J. Virol. 70: 7510-7516
- 20. Young, J.A.T. (1996) The replication cycle of HIV-1, In: The AIDS Knowledge Base. (Cohen, P., Sande, M., Volberding, P., et al., eds.) Little Brown, New York, NY in press.
- 21. Brojatsch, J., Naughton, J., Rolls, M.R., Zingler, K. and Young, J.A.T. (1996) CAR1, a TNFR-related protein is a cellular receptor for cytopathic avian leukosis and sarcoma viruses and mediates apoptosis. Cell 87: 845-855.

#### INVITED PRESENTATIONS (Meetings)

Invited Chair, Roundtable Discussion on Gene Therapies for AIDS. Second Annual NIH National AIDS Cooperative Drug Discovery and Development Meeting. California. 1988.

Invited Speaker, Banbury Conference on Receptor Mediated Virus Butry into Cells, Cold Spring Hurber Laboratory, 1991.

Invited Speaker, Keystone Symposium on Molecular Biology of Human Pathogenic Viruses. California, 1993.

Invited Speaker. Fifth Workshop on Pathogenesis by Non-acute Retroviruses. France, 1993.

Invited Speaker, Workshop on Immunology and Developmental Biology of the Chicken. Basel Institute of Immunology, Switzerland, 1994.

Invited Speaker, Sixth Workshop on Pathogenesis of Animal Retroviruses. Philadelphia 1994

Invited Speaker, Seventh Workshop on Pathogenesis of Animal Retroviruses. Seanle 1995.

Chair, Session on Receptors, Entry and Uncoating, Retrovinues Meeting at Cold Spring Harbor Laboratory, New York, 1996

Invited Speaker, FASEB Summer Conference on Principles in Viral, Bucterial, Pungal and Protozoan Pathogenesis. Colorado, 1996

Invited Speaker, FASEB Summer Conference on Virus Assembly, Vermont, 1996

Invited Symposium Speaker, American Society for Virology Symposium, Montana 1997

Invited Speaker, Cleveland Virology Symposium, 1997

Invited Speaker, Animal Cells and Viruses Gordon Conference, 1997

#### INVITED PRESENTATIONS (Institutions)

Department of Microbiology and Immunology, Penn State Medical Center, Hersbey Pennsylvania, October 1996.

Department of Mulecular Microbiology, Washington University at St. Louis School of Medicine, St. Louis, November 1996

Department of Microbiology, New York University Medical School, New York, January 1997

#### OTHER PRESENTATIONS

An attempt to specifically alter retroviral tropism using EGF-envelope chimerss. J.A.T. Young, P. Bates, II. Various. Poster presentation at Cold Spring Harbor RNA Tumor Viruses meeting, May 1988.

Transfer of susceptibility to ALSY infection into mammalism cells with chicken DNA. P. Bates, J.A.T. Young, and H.B. Varmus. Talk at Cold Spring Harbor RNA Tumor Viruses meeting. May 1989.

The human CD4 protein is efficiently incorporated into ALV particles. J.A.T. Young, P. Bates,

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#### **PUBLICATIONS**

- 1. Trowsdule, J., Young, J.A.T., Kelly, A.P., Austin, P.J., Carson, S., Meunier, H., So, A., Chrlich, H.A., Spichnan, R.S., Bodmer, J., and Bodmer, W. (1985) Structure, sequence and polymorphism in the HLA-D region. *Immunol. Rev.* 85:135-173.
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  by an HLΛ-DPα chain-specific monoclonal antibody (DPI I.1) is influenced by the
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- 8. Young, J.A.T. (1988) HIV and HLA similarity. Scientific correspondence. Nature 333:215.
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- 10. Young, J.A.T., Bates, P., Willert, K., and Yarmus, H.E. (1990) Efficient incorporation of human CD4 protein into Avian Leukosis Virus particles, Science 250:1421-1423.
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- 13. Connolly, L., Zingler, K., and Young, J.A.T. (1994) A soluble form of a receptor for subgroup A avian leukosis and sarcoma viruses (ALSV-A) blocks infection and binds directly to ALSV-A. J. Virol. 68:2760-2764.
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- 20. Young, J.A.T. (1996) The replication cycle of HIV-1. In: The AIDS Knowledge Base. (Cohen, P., Sande, M., Volberding, P., et al., eds.) Little Brown, New York, NY in pross.
- 21. Brojalsch, J., Naughton, J., Rolls. M.R., Zingler, K. and Young, J.A.T. (1996) CAR1, a TNFR-related protein is a cellular receptor for cytopathic avian leukosis and surcoma viruses and mediates apoptosis. Cell 87: 845-855.

#### INVITED PRESENTATIONS (Meetings)

Invited Chair, Roundtable Discussion on Gone Therapies for AIDS, Second Annual NIH National AIDS Cooperative Drug Discovery and Development Meeting. California, 1988.

Invited Speaker, Banbury Conference on Receptor Mediated Virus Butry into Cells, Cold Spring Hurber Laboratory, 1991.

Invited Speaker, Keystone Symposium on Molecular Biology of Human Pathogenic Viruses. California, 1993.

Invited Speaker, Fifth Workshop on Parhogenesis by Non-acute Retroviruses. France, 1993.

Invited Speaker, Workshop on Immunology and Developmental Biology of the Chicken. Basel Institute of Immunology, Switzerland, 1994.

Invited Speaker, Sixth Workshop on Puthogenesis of Animal Retroviruses. Philadelphia 1994

Invited Speaker, Seventh Workshop on Pathogenesis of Animal Retroviruses. Searcle 1995.

Chair, Session on Receptors, Entry and Uncoating, Retrovinues Meeting at Coki Spring Harbor Laboratory. New York, 1996

Invited Speaker, FASEB Summer Conference on Principles in Viral, Bacterial, Fungal and Protozoan Pathogenesis. Colorado, 1996

Invited Speaker, FASEB Summer Conference on Virus Assembly, Vermont, 1996

Invited Symposium Speaker, American Society for Virology Symposium, Montana 1997

Invited Speaker, Cleveland Virology Symposium, 1997

Invited Speaker, Animal Cells and Viruses Gordon Conference, 1997

## INVITED PRESENTATIONS (Institutions)

Department of Microbiology and Immunology, Penn State Medical Center, Hersbey Ponnsylvania, October 1996.

Department of Mulecular Microbiology, Washington University at St. Louis School of Medicine, St. Louis, November 1996

Department of Microbiology, New York University Medical School, New York, January 1997

#### OTHER PRESENTATIONS

An attempt to specifically alter retroviral tropism using EGF-envelope chimeras. J.A.T. Young, P. Bates, IL Varmus. Poster presentation at Cold Spring Harbor RNA Tumor Viruses meeting. Muy 1988.

Transfer of susceptibility to ALSY infection into mammalian cells with chicken DNA. P. Bates. J.A.T. Young, and H.B. Varmus. Talk at Cold Spring Harbor RNA Tumor Viruses meeting. May 1989.

The human CD4 protein is efficiently incorporated into ALV particles. J.A.T. Young, P. Bares,

K. Willert, and H.E. Varmun. Talk at Cold Spring Harbor RNA Tumor Viruses meeting. May 1990.

An LDL receptor-related protein is the subgroup A ALV receptor. P. Bates, J.A.T. Young, H.E. Varmus. Talk at Cold Spring Harbor RNA Tumor Viruses meeting. May 1992.

Functional characterization of the subgroup A-Avian Lenkosis Virus (ALV) receptor gene: Low levels of receptor expression are limiting for virus infection. J.A.T. Young, P. Bates. H.B. Varmus. Talk at Cold Spring Harbor RNA Tumor Viruses meeting, May 1992.

Mutational analysis of the cellular receptor for subgroup A-ALSV. K. Zingler, C. Belanger, J.A.T. Young. Talk at Cold Spring Harbor Retroviruses meeting. May 1994.

A soluble version of the subgroup A-ALSV receptor blocks infection and binds directly to ALSV-A. L. Connolly, K. Zingler, J.A.T. Young. Talk at Cold Spring Harbor Retroviruses meeting. May 1994

An assay system to determine the relative levels of intermedials and complete DNA forms of HIV-1 DNA following infection. M. Jenkins, J. Naughton, J.A.T. Young. Poster presentation at Cold Spring Harber Retroviruses meeting. May 1994

A putative receptor for cytopathic subgroups of ALSVs is a member of the Fas/TNFR protein superfamily. J. Brojatsch, J. Naughton, J.A.T. Young. Talk at Cold Spring Harbor Retroviruses meeting. May 1996.

Evidence that residue Trp-48 of TVA is involved at a step of virul entry other than binding the SU glycoprotein of subgroup A avian laukosis and sarcoma viruses. K. Zingler, J.A.T. Young. Talk at Cold Spring Harbor Retroviruses meeting. May 1996.

#### GRANTS

Characterization of ALSV-A Env/Receptor Interactions NIH: 1R29CAAI62000-01A1 \$615,301, July 1994 to June 1999

An Altempt to Turget Retrovints Vectors to Cells Expressing HIV-1 Envelope Proteins AIDS Clinical Research Center, UCSF
One-year grant (\$25,000). Funded January 26, 1994

Milion Fund (\$12,000) Harvard Medical School, July 1995

Characterizing the Mechanisms of ALSV Entry into Cells NTH: 1RO1CA70810-01 5 985, 949, July 1996 to June 2000.

#### **OUTSIDE ACTIVITIES**

1995 to present. Consultant, Chiron Corporation, Emeryville, California

1995 to present. Consultant, Vaccines and Related Biological Products Advisory

Committee, Food and Drug Administration.